

Heterogeneity of Graphite Lithiation in State-of-the-Art Cylinder-Type Li-Ion Cells

Dominik Petz,^[a, b] Martin J. Mühlbauer,^[a, c, d] Alexander Schökel,^[a, d, e] Klaus Achterhold,^[f] Franz Pfeiffer,^[f, g] Thilo Pirling,^[h] Michael Hofmann,^[a] and Anatoliy Senyshyn^{*[a]}

The two-dimensional lithium distribution in the graphite anode was non-destructively probed by spatially resolved neutron diffraction for a batch consisting of 34 different cylinder-type (18650) Li-ion batteries in fully charged state. The uniformity of the lithium distribution was quantified and correlated to the cell specifications/electrochemistry and to intrinsic cell parameters like electrode thickness, position of current collectors, etc. which were obtained by X-ray micro-computed tomography. Non-uniformities in the lithiation state of the anode from a

- [a] D. Petz, Dr. M. J. Mühlbauer, Dr. A. Schökel, Dr. M. Hofmann, Dr. A. Senyshyn Heinz Maier-Leibnitz Zentrum (MLZ) Technische Universität München Lichtenbergstr. 1, 85748 Garching, Germany E-mail: anatoliy.senyshyn@gmail.com
 [b] D. Petz
- Lehrstuhl für funktionelle Materialien, E13 Technische Universität München James-Franck-Str. 1/I, 85748 Garching b. München, Germany [C] Dr. M. J. Mühlbauer
- Helmholtz-Institute UIm for Electrochemical Energy Storage (HIU) Karlsruhe Institute of Technology (KIT)
 P.O. Box 3640. D-76021 Karlsruhe, Germany
- [d] Dr. M. J. Mühlbauer, Dr. A. Schökel Institute for Applied Materials (IAM) Karlsruhe Institute of Technology (KIT) Hermann-von-Helmholtz-Platz 1, 76344 Eggenstein-Leopoldshafen, Germany
- [e] Dr. A. Schökel Deutsches Elektronen Synchrotron (DESY) Notkestr. 85, 22607 Hamburg, Germany
- [f] Dr. K. Achterhold, Prof. F. Pfeiffer Department of Physics & Munich School of Bioengineering Technische Universität München James-Franck-Straße 1, 85748 Garching, Germany
- [g] Prof. F. Pfeiffer Department of Diagnostic and Interventional Radiology Munich School of Medicine and Klinikum rechts der Isar Ismaninger Str. 22, 81675 Munich, Germany
- [h] Dr. T. Pirling
 Institut Max von Laue Paul Langevin (ILL)
 71, Avenue des Martyrs,
 38043 Grenoble Cedex 9, France
- Supporting information for this article is available on the WWW under https://doi.org/10.1002/batt.202000178
- © 2020 The Authors. Batteries & Supercaps published by Wiley-VCH GmbH. This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

constant plateau have been observed for the majority of the studied cells. Their location corresponds to the positions of current tabs connecting the electrode stripes and areas of incomplete electrode coating at the beginning and the end of the electrode stripes. Four commonly used schemes of current lid connection were identified. Each of them displays its own effect on the uniformity of the lithiation at the anode and, therefore, variation of the intrinsic state-of-charge distribution and, most probably, the ageing behavior of the electrodes.

1. Introduction

Energy storage media based on Li-ion technology are peaking in their popularity for power and energy supply and implemented/embedded in wearable and portable electronics, cordless and communication tools and/or electric drivetrains. Such a broad range of applications for Li-ion batteries and their day-to-day use bring lifetime, cycling stability and safety issues to the same level of significance as energy and power density.^[1-3] It is well known that lithium ion batteries lose part of their capacity upon cycling and due to calendar aging, which involves different fading mechanisms, e.g. loss of movable lithium, electrolyte decomposition and degradation of active electrode material, which is supplemented by an increased internal resistance of the cell, thus reducing efficiency.^[4,5] It is worth to note that the majority of fading mechanisms accompanying standard cell operation are heterogeneous in their nature.^[6,7] This can be illustrated by a simple consideration: for example, the rate of reactions occurring in lithium-ion batteries are highly temperature dependent;^[8] cell charge/ discharge (as a source of non-uniform current density distribution)^[9-12] produces temperature gradients in different cell parts,^[13] leading to different reaction rates, and by this, to a difference in fading. The heterogeneous character of cell operation leads to a non-uniform lithium-ion distribution inside the cell,^[14,15] which affects the cell performance, lifetime and safety.

The use of battery modules equipped with thermal management systems further adds to the complexity of predicting and describing the reaction mechanisms. Furthermore, in order to reach an optimum volumetric energy density, modern lithium-ion cells are manufactured in various shapes. A clear trend towards an increase of capacity on the single cell level can be observed, where issue of cell uniformity and relevant tools for its studies gain in importance. Lithium-ion

Batteries & Supercaps 2021, 4, 327-335



batteries are closed electrochemical systems, which can easily be driven out of their thermodynamic equilibrium and change their state upon opening (by, e.g., redox reactions of electrode material, evaporation of the electrolyte, change of mechanical pressure, modified wetting conditions and electrical contacts of the electrodes and so on). Therefore, battery R&D requires nondestructive experimental techniques capable to deliver information under real operating conditions. As such, an *in situ* technique capable to unambiguously determine the lithiation grade of the electrodes without cell disassembly is desirable. Another important aspect is that the technique should be capable to deliver depth information in order to probe its spatial distribution. Among the variety of non-destructive techniques, neutron scattering (especially diffraction) is particularly suited due to numerous advantages:

- Neutrons are weakly interacting with matter in general, which leads to high achievable penetration depth and, thus, enables studies of bulky samples or objects in complex environments.
- The wavelength of thermal neutrons is comparable to interplanar spacings in crystal structure, enabling accurate neutron diffraction studies. The atomic form factor is independent from $\sin(\Theta)/\lambda$ because of the interaction with the nuclei, which results in accurate structure factors and, correspondingly, high quality structural data.
- Neutrons are not only isotope-sensitive, but also show large variation of scattering length, i.e. light elements can be localized next to heavy ones because of their distinctly different scattering cross sections. Therefore, neutron diffraction displays an enhanced sensitivity to light atoms (including lithium) and to transition metals making it complementary to X-ray diffraction, scattering and absorption. An isotope contrast can further improve the sensitivity and localization power, e.g., using isotope markers.

Neutron scattering is actively explored in battery research and, indeed, neutron diffraction is one of the most commonly used technique in the field. It offers a non-destructive way to assess the lithiation degree of anode and cathode, which is reflected by the response of the crystal structures on the momentary state-of-charge (SOC)^[16] and state-of-health (SOH).^[17] Such kind of *in situ* neutron diffraction studies are performed on Li-ion cells of different designs, either laboratory^[18–21] or commercial^[22–24] type. In recent years a number of neutron diffraction studies dedicated to the lithiation of either positive or negative electrode materials,^[25–31] aging behavior^[32,33] and effects of temperature.^[34,35]

The majority of the above-mentioned reports focuses on the structure evolution obtained by integration over a large cell volume, assuming the studied cells to be uniform. Literature on the experimental studies of lithium uniformity on the cell level is quite limited: Cai et al.^[36] reported a non-uniform degradation in large format pouch bag cells using time-of-flight neutron diffraction as a local probe; Yu et al.^[37] proposed an approach for simultaneous studies of local temperature, SOC and strain. A method for probing the 2D lithium distribution in the graphite anode using monochromatic neutron diffraction (alternative to time-of-flight neutron diffraction) was proposed in literature.^[38] The method was successfully applied to monitor the lithium distribution in four 18650-type cells adopting different configurations of current tabs^[14] and to probe the lithium distribution in variously fatigued high energy lithium cobalt oxide LCO | C^[39] and high performance lithium nickel cobalt aluminum Oxide NCA | C^[32] cells. Recently, an attempt to localize and quantify the electrolyte in frozen state was made by a similar experimental approach at low temperatures.^[33] The lithium distribution has been found strongly affected by the configuration of the current tabs and by the cell SOH leading to an increasingly heterogeneous character of cell fatigue. Besides neutron diffraction there is a number of neutron-based scattering and spectroscopic techniques capable to localize lithium (e.g., neutron depth profiling,^[40] neutron imaging,^[24] neutron reflectometry^[41] and small/wide angle neutron scattering,^[42] etc.), where neutron diffraction seems to be the optimal solution for non-destructive studies of assembled cell and spatial resolution on mm scale.

Therefore, previous studies^[14] were extended to a larger batch of cylinder cells of different type. The aim of the current work is to establish a correlation between the experimentally determined spatial distribution of lithium×in lithiated graphites Li_kC_6 , the electrochemical performance of the cell, i.e. its capacity and/or nominal currents as well as details of the inner cell organization and morphology like electrode thickness, position of current tabs, etc. In the frame of the current study a batch consisting of 34 different cells was investigated, where their cylinder type (18650) shape, rolled electrode stacks and graphite anode were common criteria for the cell selection.

Experimental Section

Galvanostatic cell cycling

The batch of 18650-type Li-ion secondary cells was composed of various types produced by different manufacturers. Their electrochemical characterization was carried out using a Neware BTS4000 multichannel potentiostat applying constant current (CC) discharge and constant current constant voltage (CCCV) charge protocols. For sake of comparison to previous data and to minimize the effects driven by relaxations and drifts on the cell parameters, the applied current in the CC part of charging/discharging was set to 400 mA and the cut-off current in CV region was set to 1% of the nominal 1 C current (different for each cell type). Cell-specific voltage windows were applied, where the lower and upper voltage boundaries were taken from the nominal voltage values provided by the cell manufacturers. An overview of the cycling parameters is given in Table 1.

X-ray computed tomography

A high-resolution phoenix v|tome|x 240s system was utilized for characterization of the cells by X-ray micro-computed tomography (X-Ray microCT). The acceleration voltage was set to 130 kV and the current to 100 μ A either using direct or transmission tube. Data collection was performed using a CCD detector, DXR-250RT General Electric, with a CsI scintillator of 700 mm thickness, an active area of 1000×1000 pixels and a pixel size of 200 μ m×200 μ m. For each cell 1001 projections (X-ray radiography images) were produced

328

	Tak	Table 1. Summarized list of parameters and obtained characteristics of studied 18650-type cells.	ist of parameters and	d obtained o	characteristi	cs of stud	led 18650-tyl	pe cells.												
$ \ \ \ \ \ \ \ \ \ \ \ \ \ $	ž		Cell type	Capacity [mAh]	Current ⁶	[A]	Voltage	Tab	Current	tab [cm		ickness ^[b]	Concentra	ation×in I	-i _x C ₆	Parame	eters pla	teau	C ^[c] excess
1 Simulting (R1 8863) 200 ⁽⁴¹⁾ 200 192.5 25.4.3 31.6 23.4.3 30.6 23.4.3 30.6 23.4.3 30.6 23.4.3 30.6 30.6.3 30.7.3 30.6.3 30.7.3 30.6.3 30.7.3 30.6.3 30.7.3 <th></th> <th></th> <th></th> <th>nominal</th> <th>observed</th> <th>charge</th> <th>discharge</th> <th>[V]</th> <th>scheme</th> <th>"+"</th> <th>-" "_"</th> <th></th> <th>m]</th> <th>plateau</th> <th>median</th> <th>mean</th> <th>μ</th> <th>X_{c}</th> <th>W</th> <th></th>				nominal	observed	charge	discharge	[V]	scheme	"+"	-" "_"		m]	plateau	median	mean	μ	X_{c}	W	
2 simular C(1860)30F 2000 249(1 3 6 275-4.3 1 540 235 041 048 088 0000 1135 5 smunary C(1860)20F 2000 249(1 2 544 2 54 23 25 037 088 088 088 008 01113 6 Simunary NR18503012FC 2000 2800.2 215 23 25 23 25 23 25 23 26 23 088 088 088 088 081 027 036 089 0011135 7 Simunary NR185032FC 2000 2800.2 25 25 25 23 23 23 25 23 23 23 25 23	-	Samsung	INR 18650 20R ^[d]	2000	1905.2	4	22	2.5-4.2	e	5.32			e	0.839	0.826	0.812	0.660	0.839	0.027	1.267
3 smrung (K1 8660 JF 200 2467, 25 5, 275-42 7 5, 54 2, 25 44 312 - 370 055 055 055 055 056 059 0105 1124 5 smrung (K1 8660 JF 200 2382, 2 13 2, 254-42 7 54 312 - 370 0567 0567 056 0591 057 056 0501 1135 6 LG (N1 81660 JR 200 2382, 3 15 2, 254-42 7 54 312 - 377 0567 056 0581 057 056 0501 1135 7 smrung (K1 8660 JF 200 2383, 4 2 0 25-42 7 5 54 312 - 377 0567 056 0581 056 057 056 000 11 135 8 smrung (K1 8660 JF 200 2383, 4 2 0 25-42 7 5 54 317 5 3 23 0 667 0567 0566 0581 056 0591 013 1 smrung (K1 8660 JF 200 2383, 4 2 0 25-42 7 5 53 310 - 279 0581 0581 0581 0581 0591 051 135 1 smrung (K1 8660 JF 200 2393) 2 15 10 2,57-42 1 5 35 2,10 0581 0681 0581 0591 0591 013 1 smrung (K1 8660 JF 200 2393) 2 15 10 2,57-42 1 5 35 2,75 - 394 081 0681 0593 0593 0593 0593 0593 0593 139 1 smrung (K1 8660 JF 200 277) 1 22 44 2 1 5 35 2,75 - 394 081 081 081 074 135 1 smrung (K1 8660 JF 200 277) 1 22 44 2 2 5,3 2,74 3 1 5 4,3 2,8 - 327 057 0581 0581 0594 001 111 1 smrung (K1 8660 JF 200 277) 1 25 4 2 2 5,3 2,74 3 1 5 4,3 2,8 - 327 0 687 0680 0593 050 014 125 1 smrung (K1 8660 JF 200 277) 1 25 4 2 2 5,3 2,74 3 1 5 4,3 2,8 - 324 0 641 050 1 smrung (K1 8660 JF 200 2377 159 145 1 smrung (K1 8660 JF 200 2377 159 145 17 12 12 12 12 12 12 12 12 12 12 12 12 12	7	Samsung	ICR 18650 30 A	3000	2694.1	ŝ	6	2.75-4.35	-	5.49	3.19 -	32	5	0.812	0.811	0.803	0.818	0.812	0.018	1.289
4 Samung NR Reso 200 205 <td>m</td> <td>Samsung</td> <td>ICR 18650 26F</td> <td>2600</td> <td>2450.7</td> <td>2.6</td> <td>5.2</td> <td>2.75-4.2</td> <td>-</td> <td>5.40</td> <td>2.85 -</td> <td>36</td> <td>5</td> <td>0.857</td> <td>0.855</td> <td>0.841</td> <td>0.648</td> <td>0.858</td> <td>0.010</td> <td>1.148</td>	m	Samsung	ICR 18650 26F	2600	2450.7	2.6	5.2	2.75-4.2	-	5.40	2.85 -	36	5	0.857	0.855	0.841	0.648	0.858	0.010	1.148
5 Smaung NR 8860 MI 3326 2 13 256-42 2 544 15 25 64 057 0897 0872 0586 0674 011 1325 7 Smuung NR 8600 X0 3000 3206 3 15 3 47 25 44 25 44 25 44 25 44 25 31 088 0887 0874 0880 0001 113 5 Smuung NR 8600 XL 1500 1201 1202 125 35 31 53 31 53 31 53 31 53 31 33 31 33 31 33 31 33 31 33 31 33 31	4	Samsung	ICR 18650 22P	2200	2062.5	2.15	10	2.75-4.2	2	5.39	2.81 -	32	9	0.900	0.887	0.872	0.558	0.900	0.019	1.125
6 Int Biology (M) 3500 25-42 1 491 35 3 50 086 057 0361 <td>Ś</td> <td>Samsung</td> <td>INR 18650 35E^[d]</td> <td>3500</td> <td>3328.6</td> <td>2</td> <td>13</td> <td>2.65-4.2</td> <td>2</td> <td>5.44</td> <td>3.12 -</td> <td>36</td> <td>2</td> <td>0.884</td> <td>0.872</td> <td>0.852</td> <td>0.566</td> <td>0.884</td> <td>0.016</td> <td>1.174</td>	Ś	Samsung	INR 18650 35E ^[d]	3500	3328.6	2	13	2.65-4.2	2	5.44	3.12 -	36	2	0.884	0.872	0.852	0.566	0.884	0.016	1.174
7 Samsung NRI N6500 3000	9	, FG	INR 18650 MJ1	3500	3021.6	3.4	10	2.5-4.2	-	4.91	3.36 -	36	2	0.867	0.860	0.841	0.577	0.867	0.011	1.135
8 Samurging R Samurging C Samurging Samurging <t< td=""><td>~</td><td>Samsung</td><td>INR 18650 30Q</td><td>3000</td><td>2896.2</td><td>m</td><td>15</td><td>2.5-4.2</td><td>с</td><td>5.40</td><td></td><td></td><td>ņ</td><td>0.894</td><td>0.883</td><td>0.865</td><td>0.573</td><td>0.896</td><td>0.018</td><td>1.122</td></t<>	~	Samsung	INR 18650 30Q	3000	2896.2	m	15	2.5-4.2	с	5.40			ņ	0.894	0.883	0.865	0.573	0.896	0.018	1.122
9 Samsung NR 1850.1E 300 1312 132 11 Samsung NR 1850.1E 500 1307 145 18 547 2 331 0.47 0.981 0.011 112 2 samsung CR 18650.26 2500 25012 2.5 5.4 2.57 2 332 0.81 0.81 0.81 0.011 113 2 samsung CR 18650.26 2500 25012 2.5 5.4 1.5 5.4 2.7 0.831 0.011 113 2 samsung CR 18650.26 3500 2575.4.2 1 5.4 2.7 3 347 0.831 0.011 113 2 samsung CR 18650.A 300 25912 2.03 0.831 0.017 1139 15 Pansonic NR 18650.A 300 23912 1.035 1.25 2.54.4.2 2.54.1 2.54 2.7 0.881 0.017 1.109 16 Pansonic NR 18650 A <td< td=""><td>∞</td><td>Samsung</td><td>INR 18650 25R</td><td>2500</td><td>2383.8</td><td>4</td><td>20</td><td>2.5-4.2</td><td>с</td><td>5.28</td><td></td><td></td><td>4</td><td>0.885</td><td>0.868</td><td>0.855</td><td>0.474</td><td>0.886</td><td>0.014</td><td>1.139</td></td<>	∞	Samsung	INR 18650 25R	2500	2383.8	4	20	2.5-4.2	с	5.28			4	0.885	0.868	0.855	0.474	0.886	0.014	1.139
10 Samsung R18 S5-4.2 1 S35 310 0311 0327 0311 0327 0311 0312 0314 0321 131 12 Samsung CR1 8650 2H 2500 2001. 2.5 4.4 2.75-4.2 1 5.3 5.5 - 340 0811 0.631 0.631 0.611 0.11 117 5amsung CR1 8650 2H 2500 2010. 2.55 3.1 0.683 0.683 0.683 0.631 0.611 0.17 17.17 15 Samsung CR1 8650 A 3000 2557.3 2.8 2.75-4.3 1<5.4	6	Samsung	INR 18650 29E	2900	2731.3	2.75	10	2.75-4.2	2	5.35	2.85 -	31	2	0.900	0.885	0.865	0.475	0.901	0.015	1.126
11 Samsung CRI 1865/05H 250 220 44 275-4.2 1 538 275 - 340 0810 077 0387 0011 1135 12 Samsung CRI 1865/05H 2000 2751 3 54 225 - 310 0887 0887 0873 0011 1135 13 Samsung CRI 1865/05 2900 3771 1935 56 275-4.12 1 544 275 311 0881 0887 0887 0871 0171 1139 15 Samsung CRI 1865/0 2900 3771 1935 54 255 251 254 2 517 254 2 310 0371 0387 0017 1135 15 Samyor NCRI 1865/0 2900 23773 157 25 253 356 548 526 0887 0387 0017 1135 17 Pansonic NCRI 1865/0 2900	10	Samsung	INR 18650 15 L	1500	1507.2	1.45	18	2.5-4.2	-	5.35	3.10 -	29	6	0.841	0.837	0.821	0.722	0.841	0.023	1.198
12 Samsung CR118650 38 3000 25/10 2 5 5 2 254,43 1 544 288 0.882 0.883 0.001 1131 13 Samsung CR118650 38 3000 25/10 3 54 284 28 0.883 0.883 0.001 1131 14 Samsung CR118650 38 3000 25/14 3 54 27 0.883 0.833 0.063 0.001 1131 15 SamyoPhransonic NCR1 8650 3300 25/34 3 2 2 2 54 2 2 54 2 2 331 1011 1173 15 SamyoPhransonic NCR1 8650 3300 29/34 1/475 59 2/54 2 533 336 0.041 0.081 0.031 0.031 1031 1132 15 SamyoPhransonic NCR1 8650 M 2000 2/54,42 2 533 516 <td< td=""><td>11</td><td>Samsung</td><td>ICR 18650 22F</td><td>2250</td><td>2202.1</td><td>2.2</td><td>4.4</td><td>2.75-4.2</td><td>-</td><td>5.38</td><td>2.75 -</td><td>34</td><td>Q</td><td>0.811</td><td>0.810</td><td>0.797</td><td>0.733</td><td>0.811</td><td>0.014</td><td>1.285</td></td<>	11	Samsung	ICR 18650 22F	2250	2202.1	2.2	4.4	2.75-4.2	-	5.38	2.75 -	34	Q	0.811	0.810	0.797	0.733	0.811	0.014	1.285
13 Samsung CRI 86503.308 3000 2751-9 3 6 275-4.35 1 5,44 2,8 6 2,75-4.35 1 5,44 2,8 6,8 0,847 0,847 0,847 0,847 0,017 1,191 1 Famsunic NCRI 16650 3500 3371.3 1,635 10 2,5-4.2 2 5,10 2,347 0,847 0,847 0,847 0,847 0,847 0,017 1,132 1 Famasonic NCRI 16650 3500 3371.3 1,435 5 2 5,44 2 5 317 2,44 2 317 0,847 0,847 0,847 0,847 0,017 1,132 1 Famasonic NCRI 16650 3300 2,347.2 1,37 2,33 3,33 0,33 0,347 0,397 0,347 0,397 0,347 0,387 0,317 1,323 1 Bassonic NCRI 16650 3300 2,347 3 3,25 2,442 2 5,433 3,637 0,417 0,593 0,337 0,417 1,323<	12	Samsung	ICR 18650 26H	2600	2610.2	2.6	5.2	2.75-4.2	-	5.30	2.73 -	33	4	0.883	0.882	0.875	0.720	0.883	0.011	1.117
14 Samung CRI 8650.28A 2800 2553 15 311 541 318 - 342 0.845 0.837 0.471 0.871	13	Samsung	ICR 18650 30B	3000	2751.9	e	9	2.75-4.35	-	5.44	2.82 -	31	-	0.863	0.862	0.858	0.843	0.863	0.020	1.143
5 Saryof*Panasonic NGR1 18650 3500 32711 3165 10 25-42 2 517 237 0.871 <th0< td=""><td>14</td><td>•.</td><td>ICR 18650 28 A</td><td>2800</td><td>2659.3</td><td>2.8</td><td>5.6</td><td>2.75-4.3</td><td>-</td><td>5.41</td><td>3.18 -</td><td>34</td><td>5</td><td>0.847</td><td>0.845</td><td>0.833</td><td>0.797</td><td>0.847</td><td>0.016</td><td>1.181</td></th0<>	14	•.	ICR 18650 28 A	2800	2659.3	2.8	5.6	2.75-4.3	-	5.41	3.18 -	34	5	0.847	0.845	0.833	0.797	0.847	0.016	1.181
16 Panasonic NCRI 18650 2000 2773 1935 5.8 2.5-4.2 2 5.10 2.32 0.838 0.833 0.833 0.836 0.741 0.838 0.002 1.212 18 Panasonic NCRI 18650 F 2.00 287.9 1.475 5.9 2.5-4.2 2 5.17 2.44 0.886 0.887 0.846 0.037 1.232 18 Panasonic NCRI 18650 FR 2.00 2.7492 1.62 5.33 3.86 5.48 0.25 0.846 0.017 1.135 16 SanyofPanasonic NCRI 18650 FR 2.00 2.743 1.75 5 2.5-4.2 2 5.33 3.35 5.48 0.886 0.017 1.159 17 SanyofPanasonic NCRI 18650 FR 2.300 2.745 1 2.5-4.22 2 5.33 3.31 0.29 0.887 0.007 1.115 16 CRI 18650 FR 2.350 2.174 4 20 2.5-4.22<	15		NCR 18650 GA	3500	3371.3	1.635	10	2.5-4.2	2		3.77 -	37	0	0.871	0.848	0.837	0.471	0.871	0.017	1.179
17 Panasonic NCR 18650 F 2900 2917 7.030 10 25.5.4.2 2 517 2.5.4 - 230 0.846 0.826 0.575 0.866 0.037 1.232 18 Fanasonic NCR 18650 BF 3300 23973 1.475 5 5.5.4.2 2 5317 234 - 400 0.856 0.851 0.856 0.551 0.884 0.037 1.59 1.59 1.51 1.55 5 2.5.4.22 2 5.33 3.85 5.48 2.86 0.881 0.851 0.832 0.031 1.159 1.50 7 2.5.4.42 3 5.5 3.33 1.5 5 5.5 3.33 5.16 2.89 0.881 0.031 1.129 1.125 1.5 2.5.4.42 3 5.16 2.89 0.875 0.881 0.031 1.125 1.47 5.05 0.311 0.17 1.195 1.17 2.05 0.346 0.517 0.881 0.0017	16	_	NCR 18650	2900	2797	1.925	5.8	2.5-4.2	2	5.10	2.50 -	32	2	0.838	0.833	0.820	0.741	0.838	0.025	1.232
18 Panasonic NRI 18650 A 3100 2397 1475 5.9 25.4.2 2 5.17 2.54 2 4.20 0.846 0.816 0.625 0.847 1.127 20 SanyoPanasonic NRI 18650 FM 3330 23492 1.655 4.875 2.5-4.2 3 5.33 3.55 1.75 5 5.33 3.55 -4.20 0.886 0.887 0.881 0.681 0.652 0.884 0.017 1.152 21 Sanyo NRI 18650 FM 2500 23703 1.75 5 2.5-4.2 3 3.55 3.35 1.6 0.875 0.881 0.815 0.881 0.915 1.32 23 LG CRT 18650 HE 2500 2174 4 20 2.5-4.2 3 5.17 24 3.07 0.961 0.637 0.031 1.122 24 CG RT8650 HH 2000 21375 0.891 0.015 0.592 0.034 0.567 0.034	17	_	NCR 18650 PF	2900	2812.7	2.030	10	2.5-4.2	2		2.40 -	32	8	0.847	0.840	0.826	0.759	0.847	0.028	1.212
19 Sanyo/Panasonic NRI 18650 BF 3350 2349.2 1,655 2,5-4.2 2 5,38 3.86 5,48 2056 0.861 0.515 0.889 0.017 1,195 20 Sanyo/Panasonic URI 8650 NX 2600 2370.3 1,75 5 2,5-4.2 2 5,38 3.86 5,48 0.875 0.861 0.861 0.017 1,195 21 Sanyo NCR 18650 HR 2500 2370.3 1,75 5 2,5-4.2 3 5,52 3:35 0.815 0.881 0.017 1,195 215 Sanyo NCR 18650 HR 2500 2175 5 2,5-4.2 3 5,52 3:35 0.817 0.018 1,172 216 CR1 18650 HR 2500 2175 3 5,52 3:35 5,17 243 0.875 0.887 0.017 1,125 226 CR1 18650 HR 2500 2175 3 5,52 3:35 5,17 259 0.877 </td <td>18</td> <td>Panasonic</td> <td>NCR 18650 A</td> <td>3100</td> <td>2987.9</td> <td>1.475</td> <td>5.9</td> <td>2.5-4.2</td> <td>2</td> <td>5.17</td> <td>2.54 -</td> <td>42</td> <td>0</td> <td>0.846</td> <td>0.836</td> <td>0.816</td> <td>0.625</td> <td>0.846</td> <td>0.017</td> <td>1.232</td>	18	Panasonic	NCR 18650 A	3100	2987.9	1.475	5.9	2.5-4.2	2	5.17	2.54 -	42	0	0.846	0.836	0.816	0.625	0.846	0.017	1.232
20 Samyo/Panasonic UR 18650 KNX 2600 2487.5 1.7 20 2.5-4.2 3 5.38 5.48 226 0.894 0.876 0.6515 0.884 0.017 1.132 21 Samyo UR 18650 KN 2600 2487.5 1.7 2.2 5.33 3.51 - 377 0.815 0.837 0.831 0.817 0.831 0.817 0.831 0.817 0.831 0.817 0.831 0.013 1.122 23 LG UR 18650 HE2 2500 2174.6 4 20 2.5-4.2 3 5.51 2.43 0.877 0.811 0.867 0.018 1.172 24 LG UR 18650 HE4 2500 2174.6 4 20 2.5-4.2 3 5.51 2.743 0.871 0.881 0.033 1.122 26 LG UR 18650 HB2 1500 1233.2 4 30 2.729 0.749 0.749 0.757 0.023 1.131	19		NCR 18650 BF	3350	3249.2	1.625	4.875	2.5-4.2	2	5.37	3.88 -	40	4	0.858	0.838	0.826	0.542	0.858	0.022	1.217
21 Sanyo UR 18650 FM 2600 2370.3 1.75 5 2.5 -4.2 2 5.33 3.51 - 337 0.881 0.881 0.881 0.881 0.881 0.035 1.284 22 53nyo NCR 18650 FL 2500 2174.6 1.62 7 2.5 -4.2 3 5.61 3.41 5.17 2.45 0.881 0.881 0.881 0.881 0.033 1132 23 LG KR 18650 HH 3200 313.6 1 2.5 -4.2 3 5.60 3.41 5.17 245 0.877 0.861 0.846 0.57 0.878 0.033 1132 24 LG KR 18650 HE 2500 322.6 3.1 10 2.5 -4.2 3 5.60 3.41 5.17 249 0.877 0.861 0.875 0.871 0.13 1172 25 LG KR 18650 HE 2500 1407.6 4 30 2.52 0.341 0.729 0.740 0.574 0.297 0.033 1131 26 KR 18650 HE <	20	• ·	UR 18650 NSX	2600	2487.5	1.7	20	2.5-4.2	с	5.38			9	0.894	0.876	0.861	0.515	0.894	0.017	1.159
22 Sanyo NCR 18650 BL 3350 3136.6 1.62 7 2.5-4.2 2 5.34 3.87 - 397 0.817 0.813 0.015 1.312 23 LG CR 18650 HE2 2500 2174.6 4 20 2.5-4.2 3 5.516 2.28 0.877 0.841 0.817 0.817 0.841 0.116 1.172 24 LG CR 18650 HE4 2500 2.2174.6 4 20 2.5-4.2 3 5.16 2.28 0.877 0.841 0.191 0.875 0.014 1.125 25 LG INR 18650 HB6 1500 1253.2 4 30 2.0-4.2 3 5.46 3.37 0.734 0.59 0.387 0.014 1.125 26 LG INR 18650 HB6 1500 1253.2 4 30 2.0-4.2 3 5.46 3.37 0.734 0.59 0.734 0.59 0.731 1.43 27 LG IR8650 HB2 1500 125.4.2 3 5.41 3.35 5.17 239	21	Sanyo	UR 18650 FM	2600	2370.3	1.75	5	2.5-4.2	2	5.33	3.51 -	35	2	0.834	0.826	0.815	0.682	0.832	0.035	1.284
23 LG CR 18650 HE2 2500 2174.6 4 20 25-42 3 5.516 228 0.875 0.871 0.841 0.410 0.876 0.018 1.172 24 LG IKR 18650 HE4 2500 2174.6 4 20 2.5-42 3 5.60 3.41 5.17 2.45 0.877 0.861 0.846 0.033 1.152 26 IKR 18650 HB6 1500 12372.2 4 30 2.5-42 3 5.48 3.35 - 343 0.875 0.871 0.876 0.018 1.172 26 IKR 18650 HB2 1500 1407.6 4 30 2.5-42 3 5.48 3.35 1.1 0.860 0.748 0.875 0.035 1.419 27 LG IKR 18650 HB2 1500 1407.6 4 30 2.5-42 3 5.49 3.35 5.10 2.73 0.749 0.774 0.59 0.025 0.141 28 507 US 18650 HB2 110 1175 3 5.48 3.33	22	Sanyo	NCR 18650 BL	3350	3136.6	1.62	7	2.5-4.2	2	5.34	3.87 -	39	7	0.837	0.813	0.799	0.386	0.838	0.015	1.312
24 IG ICR 18650 HE4 2500 2287.5 4 20 2.5-4.2 3 5.60 3.41 5.17 245 0.877 0.846 0.627 0.878 0.033 1.152 25 LG INR 18650 MH1 3200 3202.6 3.1 10 2.5-4.2 3 5.60 3.41 5.17 245 0.877 0.846 0.627 0.875 0.877 0.034 1.126 26 ICR 18650 HB6 1500 1202 2.5-4.2 3 5.48 3.28 5.10 2724 0.574 0.579 0.037 1.511 27 LG ICR 18650 HB 1500 12052 2.15 3.225 2.5-4.2 3 5.51 2.73 0.897 0.897 0.573 0.597 0.014 1.75 28 LG ICR 18650 BS 2.00 2109 4 25 2.5-4.2 3 5.61 3.33 5.17 239 0.887 0.849 0.677 0.592 0.014 1.156 29 ICR 18650 BS 3.40 0.57 0.868	23		ICR 18650 HE2	2500	2174.6	4	20	2.5-4.2	3	5.52			8	0.875	0.851	0.841	0.410	0.876	0.018	1.172
25 LG INR 18650 MH1 3200 3022.6 3.1 10 2.5-4.2 1 4.96 3.35 - 343 0.875 0.871 0.860 0.748 0.875 0.014 1.126 26 LG INR 18650 HB6 1500 1253.2 4 30 2.5-4.2 3 5.48 3.35 5.10 272 0.734 0.549 0.745 0.025 1.479 27 LG KR 18650 HB2 1500 1407.6 4 30 2.5-4.2 3 5.5 0.743 0.743 0.512 0.895 0.025 1.144 28 LG KR 18650 HD2 2.000 2109 4 25 2.5-4.2 3 5.68 2.77 0.865 0.817 0.512 0.895 0.011 1.455 29 Sony U5 18650 VFG 2600 210.3 1.17 2.85 3.17 3.95 0.877 0.865 0.011 1.455 30 Sony U5 18650 VFG 200 261 2.77 0.865 0.871 0.867 0.867	24	_	ICR 18650 HE4	2500	2287.5	4	20	2.5-4.2	с	5.60			5	0.877	0.861	0.846	0.627	0.878	0.033	1.152
26 LG INR 18650 HB6 1500 1253.2 4 30 2.0-4.2 3 5.48 3.28 5.10 272 0.754 0.734 0.549 0.757 0.022 1.479 27 LG ICR 18650 HB2 1500 1407.6 4 30 2.5-4.2 3 5.49 3.33 5.17 239 0.897 0.817 0.512 0.895 0.022 1.174 28 LG ICR 18650 HB2 1500 1407.6 4 30 2.5-4.2 3 5.61 3.33 5.17 239 0.887 0.847 0.573 0.023 1.151 29 LG ICR 18650 HB2 1500 261.3 4 30 2.5-4.2 3 5.61 3.33 5.17 2.39 0.877 0.873 0.871 0.871 0.871 0.871 0.871 0.871 0.871 0.871 0.871 0.871 0.871 0.871 0.871 0.872 0.011 1.45 30 30 30 30 0.911 1.76 33 2.413 3.73 0.885 <td>25</td> <td>_</td> <td>INR 18650 MH1</td> <td>3200</td> <td>3022.6</td> <td>3.1</td> <td>10</td> <td>2.5-4.2</td> <td>-</td> <td>4.96</td> <td>3.35 -</td> <td>34</td> <td>ņ</td> <td>0.875</td> <td>0.871</td> <td>0.860</td> <td>0.748</td> <td>0.875</td> <td>0.014</td> <td>1.126</td>	25	_	INR 18650 MH1	3200	3022.6	3.1	10	2.5-4.2	-	4.96	3.35 -	34	ņ	0.875	0.871	0.860	0.748	0.875	0.014	1.126
27 LG ICR 18650 HB2 1500 1407.6 4 30 2.5-4.2 3 5.49 3.38 5.20 265 0.743 0.724 0.529 0.745 0.023 1.11 28 LG ICR 18650 S3 2200 2205 2.15 3.225 2.75-4.2 1 4.90 3.31 - 391 0.895 0.871 0.512 0.895 0.023 1.14 29 LG ICR 18650 H2 2.000 2109.8 4 25 2.5-4.2 3 5.61 3.33 5.17 239 0.897 0.844 0.571 0.865 0.0101 1.175 30 Sony US 18650 VTC5 2600 2612.3 4 30 2.0-32 2.54 2.3 5.68 2.71 0.865 0.871 0.871 0.872 0.011 1.175 31 Samsung ICR 18650 B 3.400 3202 1.1 2.2 2.54.2 3.14 345 0.871 0.572 0.690 0.871 0.784 0.571 0.191 1.156 367 368	26	_	INR 18650 HB6	1500	1253.2	4	30	2.0-4.2	с	5.48			2	0.754	0.740	0.734	0.549	0.757	0.025	1.479
28 LG ICR 18650 S3 2200 2205 2.15 3.225 2.75-4.2 1 4.90 3.31 - 391 0.895 0.865 0.817 0.512 0.895 0.023 1.151 29 LG ICR 18650 HDz C 2000 2109.8 4 25 2.5-4.2 3 5.561 3.33 5.17 239 0.890 0.871 0.844 0.677 0.865 0.031 1.175 30 Sony US 18650 VTG5 2600 2612.3 4 30 2.0-31 1.1 2.2 2.5-4.2 3 5.68 2.17 239 0.895 0.871 0.874 0.677 0.671 0.872 0.011 1.175 31 Samsung ICR 18650 L6H 2200 2202 1.1 2.2 3.0-4.2 1 5.42 3.14 345 0.676 0.871 0.872 0.011 1.175 31 Samsung ICR 18650 L6F 2.00 2.324 1.3 2.54 2.33 2.57 4.13 0.676 0.670 0.872 0.011 1.	27	ГG	ICR 18650 HB2	1500	1407.6	4	30	2.5-4.2	с	5.49			5	0.743	0.729	0.724	0.529	0.745	0.022	1.511
29 LG ICR 18650 HD2 C 2000 2109.8 4 25 2.5-4.2 3 3.3 5.17 239 0.890 0.871 0.884 0.573 0.890 0.023 1.151 30 Sony US 18650 VTC5 2600 2612.3 4 30 2.0-4.2 3 5.68 3.29 5.68 271 0.865 0.877 0.844 0.677 0.815 0.11 1.445 31 Samsung ICR 18650 26H 2200 2327 1.1 2.2 5.24 3.14 345 0.777 0.764 0.677 0.872 0.011 1.445 32 Samsung ICR 18650 26F 2200 2467.6 1.3 2.6 3.0-4.2 1 5.49 3.17 345 0.772 0.670 0.872 0.014 1.267 33 Samsung ICR 18650 26F 2600 2467.6 1.3 2.6 3.0-4.2 1 5.49 3.17 345 0.872 0.690 0.617 1.199 33 Samsung ICR 18650 26F 2600 2.13	28	_	ICR 18650 S3	2200	2205	2.15	3.225	2.75-4.2	-	4.90	3.31 -	39	-	0.895	0.865	0.817	0.512	0.895	0.025	1.144
30 Sony US 18650 VTC5 2600 2612.3 4 30 2.0-4.2 3 5.68 3.27 0.865 0.857 0.844 0.677 0.865 0.019 1.175 31 Samsung ICR 18650 26H 2200 2237 1.1 2.2 3.0-4.2 1 5.42 3,14 345 0.777 0.764 0.677 0.872 0.011 1.445 32 Panasonic NCR 18650 3400 3202 1.7 6.8 2.5-4.2 2 5.49 3.17 345 0.872 0.841 0.677 0.872 0.011 1.445 33 Samsung ICR 18650 26H 2.00 2467.6 1.3 2.6 3.0-4.2 1 5.49 3.17 345 0.872 0.871 1.979 1.999 34 A123 APR 18650M—A1 1100 1014 5 20 2.050 0.569 0.592 0.690 0.616 1.590 34 A123 APR 18650M—A1 1100 1014 5 20 2.0-32 0.555 305<	29	_	ICR 18650 HD2 C	2000	2109.8	4	25	2.5-4.2	Э	5.61			6	0.890	0.871	0.854	0.533	0.890	0.023	1.151
31 Samsung ICR 18650 26H 2200 2237 1.1 2.2 3.0-4.2 1 5.42 3,14 345 0.777 0.764 0.670 0.782 0.011 1.445 32 Panasonic NCR 18650B 3400 3202 1.7 6.8 2.5-4.2 2 5.23 2.57 413 0.872 0.841 0.571 0.872 0.014 1.267 33 Samsung ICR 18650B 3400 3202 1.3 2.6 3.0-4.2 1 5.49 3.17 345 0.872 0.872 0.872 0.017 1.199 33 Samsung ICR 18650B-A1 1100 1014 5 20 2.0-3.6 4 0.62 0.590 0.669 0.782 0.017 1.199 34 A123 APR 18650M-A1 1100 1014 5 20 2.0-3.6 4 0.62 0.590 0.669 0.782 0.016 1.590 34 A123 APR 18650M-A1 1100 1014 5 20 2.060 0.669 0.565 0.59	30		US 18650 VTC5	2600	2612.3	4	30	2.0-4.2	3	5.68			Ļ	0.865	0.857	0.844	0.677	0.865	0.019	1.175
32 Panasonic NCR 18650B 3400 3202 1.7 6.8 2.5-4.2 2 5.23 2.57 413 0.872 0.841 0.571 0.872 0.014 1.267 33 Samsung ICR 18650 26 2.467.6 1.3 2.6 3.0-4.2 1 5.49 3.17 345 0.872 0.879 0.872 0.017 1.199 34 A123 APR 18650M—A1 1100 1014 5 20 2.0-3.6 4 0.62 0.55 0.690 0.665 0.592 0.016 1.590 34 A123 APR 18650M—A1 1100 1014 5 20 2.0-3.6 4 0.62 0.55 0.592 0.690 0.665 0.592 0.690 0.616 1.590 34 Alzianum recommended current from the manufacturer. Note: applied charge/discharge current was 0.4 A (CCCV). [b] Thickness of full stack (copper foil - negative electrode - separator - positive electrode - s	31		ICR 18650 26H	2200	2237	1.1	2.2	3.0-4.2	-	5.42	3,14	34	5	0.782	0.777	0.764	0.670	0.782	0.011	1.445
33 Samsung ICR 18650 26F 2600 2467.6 1.3 2.6 3.0-4.2 1 5.49 3.17 345 0.872 0.879 0.785 0.872 0.017 1.199 34 A123 APR 18650M–A1 1100 1014 5 20 2.0-3.6 4 0.62 0.55 305 0.660 0.665 0.592 0.690 0.016 1.590 [a] Maximum recommended current from the manufacturer. Note: applied charge/discharge current was 0.4 A (CCCV). [b] Thickness of full stack (copper foil - negative electrode - separator - positive electrode - separator - negative electrode at SALSA diffractometer. Spatially-resolved neutron diffraction data were collected at SALSA diffractometer.	32	_	NCR 18650B	3400	3202	1.7	6.8	2.5-4.2	2	5.23	2.57	41	3	0.872	0.862	0.841	0.571	0.872	0.014	1.267
34 A123 APR 18650M—A1 1100 1014 5 20 2.0–3.6 4 0.62 0.55 305 0.690 0.665 0.592 0.690 0.016 1.590 [a] Maximum recommended current from the manufacturer. Note: applied charge/discharge current was 0.4 A (CCCV). [b] Thickness of full stack (copper foil - negative electrode - separator - positive electrode - lectrode - separator - negative electrode. [c] The graphite excess was estimated using observed cell capacity (column 5) and "mean" lithium concentration in the anode (column 15). [d] Spatially-resolved neutron diffraction data were collected at SALSA diffractometer.	33	Samsung	ICR 18650 26F	2600	2467.6	1.3	2.6	3.0-4.2	-	5.49	3.17	34	5	0.872	0.870	0.859	0.785	0.872	0.017	1.199
[a] Maximum recommended current from the manufacturer. Note: applied charge/discharge current was 0.4 A (CCCV). [b] Thickness of full stack (copper foil – negative electrode – separator – positive electrode – sum aluminum – positive electrode – separator – positive electrode). [c] The graphite excess was estimated using observed cell capacity (column 5) and "mean" lithium concentration in the anode (column 15). [d] Spatially-resolved neutron diffraction data were collected at SALSA diffractometer.	34	A123	APR 18650M-A1	1100	1014	5	20	2.0–3.6	4	0.62	0.55	30	5	0.690	0.680	0.665	0.592	0.690	0.016	1.590
auminum - postive electione - separator - regarive electronecy, ici, me graphine excess was estimated using observed centratory (communicy) and mean infinum concentration in the anore (commin 12), fully resolved neutron diffraction data were collected at SALSA diffractometer.	[a]	Maximum recomme	nded current from t	he manufac	turer. Note:	applied cl	harge/discha	rge current v	vas 0.4 A (0	CCV). [k	J Thickn	ess of fu	ull stack (co	pper foil –	negative	electrod	e – sepa	the and	bositive (electrode –
Junuary resolved region data were concreted at JALAN dimactorizete.		mmum – positive er Miallyr-rasolyad nantr	ectrode - separator	– וופטמוועפ אישים בחווסכו	electroue). I	A diffracto	apriite exces:	was esulfid	neu using o	noser vec	ו רבוו רמל	JACILY (C		un mean		OUCEULIS			ne (colul	[b] .(c1 1111
	ň	מוומווא-ובסטואבת וובתנו	טון טווומרווטון טמנמ	אבוב רחוברו	יבח מו סארסי	א מווומרור	ווופרבוי													





during full cell rotation over 360°, corresponding to an angular step of 0.36° between two projections. A single projection was averaged over three single exposures of 2000 ms each. Data reconstruction was performed using phoenix datos | x software. The analysis of the reconstructed stacks was carried out using ImageJ.^[43]

Spatially resolved neutron diffraction: experimental setup

Spatially-resolved neutron powder diffraction data on the selected Li-ion batteries were collected at the materials science diffractometer $\mathsf{STRESS}\text{-}\mathsf{SPEC}^{^{[44]}}$ at the research reactor FRM II (Heinz Maier-Leibnitz Zentrum, Garching, Germany) and the engineering diffractometer SALSA^[45] (Institut Laue- Langevin, Grenoble, France). For the spatially - resolved neutron diffraction experiments, the diffraction signal was collected from the scattering ("gauge") volume defined by neutron optics, where the dimension of the volume is typically much smaller than the size of the studied object. By moving the sample different regions can be irradiated and finally a spatially resolved distribution of structural properties can be reconstructed. A photo and a sketch of the utilized experimental setup is shown in Figure 1a-b, where the incoming, diffracted and transmitted neutron beams, which define the gauge volume, are depicted. The monochromatic neutron beam with a wavelength of $\lambda \approx$ 1.6 Å is limited to 2 mm in width and 17 mm in height in front of the sample using a slit system. In all cases, the data collection was performed at the center part of the battery in a plane perpendicular to its cylinder axis. The active volume of the studied cells can be approximated by two concentric cylinders, where the first one corresponds to the cell housing and the second one to the center pin. All relevant battery components like electrode coating materials, current collectors and separators are approximated to form an isotropic medium delivering a diffraction signal at a desired angle 2θ . A radial collimator with an acceptance width of 2 mm selects the scattered signal and defines the horizontal cross section of the gauge volume. The achieved distribution of the gauge volumes (points at which diffraction patterns were collected) is depicted in Figure 1b. A 2D position sensitive neutron detector (³He multiwire proportional chamber) with an active area of 256×256 mm² was used for data collection at a sample-to-detector distance of 1056 mm located at a central scattering angle of 26 deg 2θ . The exposure times per diffraction pattern (per gauge volume) were varied within several minutes depending on the depth of the gauge volume location. After data

corrections for detector nonlinearities, geometrical aberrations and curvature of diffraction rings, the data was integrated into normal 1D diffraction patterns (intensity vs. 2θ angle, Figure 1c). The utilized experimental setup was very similar at STRESS-SPEC and SALSA. The only difference was: for positioning of the cell with respect to the incoming and the diffracted beam, linear and rotational stages were used at STRESS-SPEC, whereas at SALSA a hexapod (Stewart platform) was utilized. An effectively smaller gauge volume was achieved at SALSA due to a radial oscillating collimator with 1 mm horizontal field-of-view. The detector at STRESS-SPEC had higher $2\theta \times \nu$ coverage ($256 \times 256 \text{ mm}^2$) compared to SALSA ($80 \times 80 \text{ mm}^2$).

Spatially resolved neutron diffraction: data evaluation

The central angle of the flat detector at $2\theta = 26^{\circ}$ had been chosen to collect the evolution of the diffraction signal from two lithiated carbons – stage I (LiC_6) and stage II (LiC_{12}), where corresponding 001 and 002 reflections are typically present in charged batteries. Crystal structure of stage I and stage II responds weakly on lithiation^[46] so that their structure factors and cell volumes can be assumed SOC independent. The relative fraction of the two phases can be used to determine the lithiation \times in Li_xC₆, which has been found extremely sensitive to state-of-charge, state-of-health and temperature and, thus, is often used for monitoring the cell performance.^[15-17,25,47] The 001 and 002 reflections for LiC_6 and LiC_{12} are well separated in 2θ , characterized by high intensity and are well suited for the calculation of the relative phases of stage I/stage II. Molar quantities were calculated as $n = I_{hkl} [V/F_{hkl}^2] [\rho/M]$, where I_{bkl} represents the observed intensity of the Bragg reflection with Miller indices *hkl*, F_{hkl} is the corresponding structure factor, V is the unit cell volume, M is the molar mass and ρ corresponds to the density. The molar phase ratio $y = n_{LiC_6}/(n_{LiC_6} + n_{LiC_{12}})$ can be transformed to a relative lithium concentration×of the lithiated graphite anode $\text{Li}_x \text{C}_6$ with x = 1/(2 - y). Thus \times yields 1 for pure stage I with y = 1 and x = 0.5 for pure stage II and it can be used as a measure for the SOC. Typically the 001 reflection for LiC_6 vanishes at SOC < 50%, so that \times in Li_xC₆ cannot be unambiguously determined. For the current study of cells in fully charged state at least traces of the LiC₆ 001 reflection were present throughout all collected datasets. No signal from low lithiated graphites ($LiC_{x>12}$) were noticed for the studied cell series with the given instrumental peak and spatial resolution.

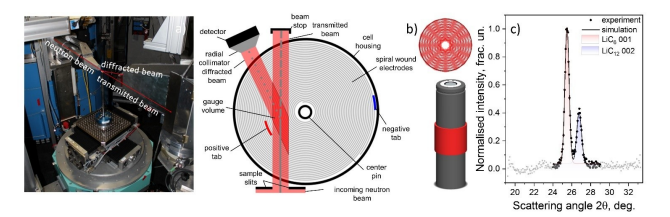


Figure 1. Photo of experimental setup (a); scheme of neutron flight path of incoming, diffracted and transmitted beam in the spatially-resolved diffraction experiment (b) along with the sketch of studied gauge volumes and the scanned area; example diffraction pattern taken from the gauge volume showing LiC_6 001 and LiC_{12} 002 reflections (c).



2. Results and Discussion

Relevant cell parameters are listed in Table 1. In the current study the cells were cycled at 400 mA, which is much smaller than the nominal charge/discharge current recommended by the cell manufacturers. Nevertheless, despite the relatively "mild" cycling conditions, the obtained cell capacities were generally less than the nominal capacities specified by the cell manufacturers by 5% in average. No clear correlation between capacity differences and other cell parameters could be found on first glance.

From Table 1 it can be seen that the cell capacity displays a complex correlation character to the voltage window, charge/ discharge currents, thickness of electrodes, etc. The correlations can hardly be drawn by direct comparison, but become evident when the electrical transport in the cells is included into consideration. As mentioned in Ref. [14], the current tabs/lids play a crucial role for the cycling performance of the battery. The position of the electrical connection to positive and negative current collectors defines the shape of the current distribution along the electrode stripe and determines the maximum allowed cycling (discharge) current. Temperature is another factor defining the magnitude of the current: the area where a current tab is electrically connected to a current collector becomes significantly heated due to resistive current losses (Joule heating), in comparison to the rest of the cell. Therefore, applications of extensively high discharge currents can hardly be achieved using a single current tab at the negative current collector and, therefore, two or more current collectors are often required on the anode side. It is worth to note that besides the electrode length the current density does also affect the lithium distribution over the thickness either on positive^[48] or negative^[49] electrodes. When no current is applied the instantaneous lithium distributions are relaxed to an intermediate lithium concentration, where time constants depend on the applied current density.^[50]

The tab configurations were extracted non-destructively using X-ray microCT, where the number of current tabs, their dimensions and positions in the stack were determined. Four connection schemes were observed utilizing either two – or three – tab connections to the current collectors¹. In cells with three-tab connections two negative ("–") current tabs to the anode were present against one positive ("+") current tab at the cathode. In all studied cells at least one "– " current tab was contacted at the outer end of anode stripe and to the cell housing. In cells adopting three-tabs another "– " current tab was placed at the opposite end of the anode stripe and the "+" current tab was located at the middle of the electrode stripe (Scheme 3). In the two-tab schemes the "+" tab was observed either at the center pin connecting the cathode stripe at its

opposite (inner) end (Scheme 1) or at the middle of the stripe (Scheme 2). As mentioned in Ref. [14], the connection of the electrode stripes via Scheme 1 usually produces a more uniform current distribution than Scheme 2 with the current supply at the middle. Interestingly, no preferences to one or another connection scheme was noticed (Scheme 2 was realized in 11 out of 34 cells; Scheme 1 in 12 out 34). Cell 34, a high power LiFePO₄-based cell from A123, is characterized by a unique connection configuration (Scheme 4), where the positive and the negative current tabs both are connected in the middle of the positive and the negative current collectors/ electrodes² leading to a configuration potentially providing the lowest internal resistance.

The surface uniformity is affected by the current tabs. Besides the number and the position of the current tabs, their length is another crucial parameter defining the maximum current flow and the out-of-plane (vertical) uniformity of a 18650-type cell. Analysis of the current tab dimensions yielded a few empirical relationships:

- Cells suited for nominal discharge current above 15 A have two current tabs at the negative electrode;
- Length of "positive" current tab is linearly proportional to the applied currents;
- All studied cells can be split into four groups adopting a specific relationship between the length of the positive and the negative current tabs.

Indeed, X-Ray microCT revealed, that current tabs at the positive electrode have a length typically varying from 4.90 to 5.68 mm. In general, current tabs at the negative electrode (either single or double) are shorter³ and their length is proportional to the ones at the opposite electrode (see Figure S1-S2). The analysis of length ratios of the current tabs displays several discrete values, which are characteristic to the cell manufacturer and the number of current collectors in the cell. There is a certain relationship between the cell capacity and the way how the current collectors are connected (Figure S3): in accordance with Table 1 the 18650-type cells built on Scheme 1 are suited for low power applications and are characterized by capacities in the range of 2000-3000 mAh; highest cell capacities are achieved applying Scheme 2; Schemes 3 and 4 are justified for cells in high power applications.

Another important parameter is the electrode thickness, which tunes the cell-specific C-rate capability and capacity. Thicknesses of the complete electrode stack including separator, negative and positive battery electrode coating and current collectors are listed in Table 1; thickness vs. discharge C-rate dependence is plotted in Figure S4. It can be seen that the electrode thickness in high power 18650-type cells (built either via Scheme 3 or 4) does not exceed 300 μ m. For cells with electrode stack is spread from 300 to 390 μ m, whereas the thicknest electrode stacks (up to 420 μ m) are found in cells using Scheme 2, which leads to the highest cell capacities.

The lithium concentrations obtained by spatially-resolved neutron powder diffraction are depicted in Figure 2 by false colors for all studied 18650-type cells. All data were collected at

¹Ten out of 34 studied cells had three current tabs.

²Current tabs utilized in Scheme 4 are short and, therefore, not visible in the collected diffraction data of the middle of the cell height.

³Except cell 20, where the current tab at the negative collector has been found longer and cells 8 and 30 exhibiting nearly equally long current tabs at positive and negative current collectors.



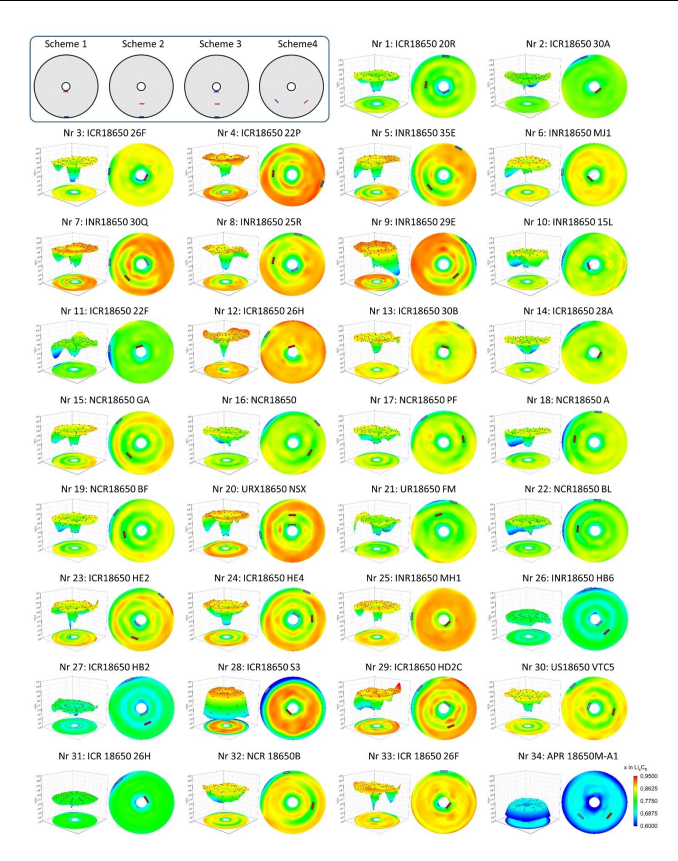


Figure 2. Lithium concentration in the graphite anode of different 18650-type cells. Data were taken at middle height. Lithium concentrations are presented by false colors, color scheme and scales are chosen to be the same for all cells. Negative and positive current tabs are presented by the black bars inside the concentration plots; blue and red colors correspond to anode and cathode current tabs. Schematic representation of electrode connections according to Schemes 1–4 is represented at the upper left corner.

Batteries & Supercaps 2021, 4, 327-335

332

 $\ensuremath{\mathbb C}$ 2020 The Authors. Batteries & Supercaps published by Wiley-VCH GmbH



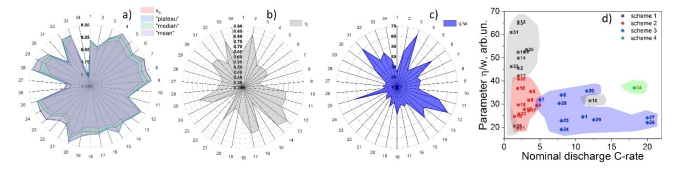


Figure 3. Spider chart diagrams of statistical parameters defining the lithium distribution (a), amount of the constant concentration plateau in the distribution histogram (Figure S5) (b), distribution of a plateau-quality parameter η/w over the studied 18650-type cells (c) and its correlation to discharge C-rate and electrode connection Schemes 1–4 (d).

the center height of the 18650-type cell. The anode and the cathode current tabs, obtained from X-ray CT, are depicted by black bars in the concentration plots with blue and red marks (corresponding to negative and positive electrodes) in the cell schematics. The obtained 2D distributions of lithiation levels have been found very different in shape and magnitude. The studied cells are suited for different applications and therefore adopt different cell chemistries and are specified for a variety of cycling currents. Different cell balancing along with an individual initial capacity loss during cell formation leads to a non-trivial correlation between cell capacity and lithiation of the anode⁴.

Each cell displays its unique lithium distribution, which is characterized by a certain degree of heterogeneity, nevertheless several systematic observations can be listed:

- (I) at places, where current tabs are attached to electrodes, clearly reduced graphite lithiation is observed, which can be attributed to a lack of electrode coating at these positions. Similar regions occur at the beginning and the end of each electrode stripe due to missing electrode coatings.^[32]
- (II) cells with electrode connections according to Scheme 1 and 2 display the most uniform lithium distribution characterized by a clearly defined 2D plateau behavior;
- (III) lithium non-uniformities at the center of the electrode stripes are observed for cells with electrode connection of Scheme 2 and 3, where the area of the observed nonuniformities was quite different.

Observation (III) is to a large extent affected by the spatial resolution of the experimental setup. Nevertheless, the presence of rather "narrow" non-uniformities in the middle of the electrode roll of cells 4–5 to highly non-uniform cases (ring shaped) as in cells 15, 19, 26 or 27, point to the real character of heterogeneous distribution in these cells. Furthermore, statement (I) has to be extended by considering the temperature: near the current tabs the highest current density and,

correspondingly, the highest temperature is supposed to occur being a potential reason for non-uniformity.

A more detailed analysis of the 2D lithium distributions presented in Figure 2 requires quantitative indicators capable to characterize the properties of the distribution, its uniformity and conformity. In Ref. [32], several statistical parameters were proposed to characterize the observed 2D distribution, namely the mean value and its standard deviation describing the average lithium concentration $\langle x_{Li} \rangle$, the median of the distribution and the plateau value x_p . As recently shown, these statistic indicators perform well for a set of variously aged cells belonging to a common cell type and the same batch. However, practice shows that "mean", "median" and "plateau" parameters obtained for different cell types cannot be correlated directly to other cell properties listed in Table 1 due to different chemistries, balancing and irreversible capacity loss upon cell formation. A direct comparison, consequently, requires a more generalized approach. For example, instead of using integral statistical parameters, the uniformity of the distribution can adequately be quantified by the analysis of distribution histograms. The histograms of the lithium distributions displayed in Figure 2 are shown in Figure S5. Their shape is significantly different from a δ -function expected for an ideal uniform system⁵. Instead of that it can be represented by a main peak corresponding to the constant lithiation plateau and a broad hump caused by non-uniformity. The plateau peak was fitted with a gaussian to obtain the estimate for $\!\times\!in$ Li_xC_6 for the plateau value corresponding to the peak position x_c , and the homogeneity of the plateau – width of the peak w. Degree of homogeneity corresponding to the amount of plateau area in the whole histogram was calculated as $\eta = A_p/A_c$, where A_p is the integral intensity of the gauss fit and $A_c = \int histogram$ is the integral over the whole histogram.

The x_c parameters were obtained for each cell (see Table 1) and are plotted in Figure 3a together with "mean" and "median" indicators in the form of a "spider" chart. All three indicators are very close with the best agreement between x_c and the "median" values. Besides x_c and w describing the constant concentration plateau and its "width", the integral uniformity parameter η was estimated for each cell and are plotted in Figure 3b. A list of cells (Nr. 13, 2, 14, 33, 17) is yield where a uniform plateau-like behavior is dominating. Accord-

⁴Excess of carbon used for cell balancing purposes was determined on the basis of observed cell capacities and experimental lithium concentrations in the graphite and is listed for each cell in Table 1 (column 20).

⁵An isotropic lithiation level × in Li_xC₆ for the entire anode is expected to reach an equilibrium state after charging with extended CV with low current cut-off values, e.g., $l_c < Q/100$ or less

European Chemical Societies Publishing

ingly, evaluation of "1- η " (Figure S6a) creates a list of cells with the most heterogeneous lithiation levels (22, 23, 15, 8, 9), where no obvious correlation to cell capacity, nominal discharge current, electrode thickness etc. (Figure S6) can be established.

Even in cells with a dominant plateau region the lithiation plateau is often not ideally flat (constant), i.e. its perfectness is determined by the width w_1 (Figure S6c) defining its deviation from the constant. Again, several cells with relatively high w can be identified (Nr. 21, 24, 17, 1, 16) along with the list of cells (3, 6, 12, 31, 25) where deviations of the plateau values are minimal. Based on this, a plateau-quality parameter η/w (Figure 3c) can be proposed⁶, which is an area fraction of the plateau with respect to its "width". The best plateau-quality parameters were observed for cells 12, 3, 31, 6, 25.

A closer look on the cell properties revealed a clear correlation between plateau-quality parameter, location of the lids and discharge current (Figure 3d). Although the rather qualitative fitting approach, the results in general have been found in good agreement with Figure 2. One can notice that cells built according to Scheme 1 are characterized by the best defined plateau of constant lithium concentration. Electrode connection schemes 2–4 generally lead to a more heterogeneous behavior resulting in a non-uniform spread and less pronounced concentration plateaus, which might be an intentional compromise maintaining internal cell resistance low.

3. Conclusions

A number of 34 Li-ion cells of 18650-type from different manufacturers was studied. The chosen cells were based on different cell chemistries and were designed for different applications, charge currents, protocols and voltage windows. Common features of the studied cells are their cylinder type shape, rolled electrode stacks and graphite anode, which were the factors defining the cell selection. The distribution of lithium in the graphite anode for the set of Li-ion batteries was probed using a combination of spatially-resolved neutron diffraction, X-ray computed tomography and electrochemistry. Studies were performed in fully charged state using low cycling currents and CCCV charging protocol with small cut-off⁷.

All studies were performed non-destructively. The cell layout: thickness of electrode stacks and details of its winding, position, length and the number of current tabs, center pins, gas vent and electric contacts were determined for each cell. Correlations between the number, length and position of current tabs, thickness of electrode stack, cell capacity and nominal cycling currents (C-rates) were established.

The 2D lithium distribution \times in Li_xC₆ was extracted from diffraction intensities at the middle height of each cell. The location of the current tabs and their connection to the

electrode stripes have been found correlated to the lithium distribution inside the graphite anode. Four commonly used schemes of the current lid connection were identified, where cells with an electrode connection in line with Scheme 1 display the most uniform lithiation; electrode connections via schemes 2 and 3 lead to lithium non-uniformities at the center of the electrode stripes: the cell build on Scheme 4 is characterized by the largest non-uniformity with a very narrow concentration plateau and a vertical gradient of lithium concentrations.^[14]

The performed experiments suffered from the limited spatial resolution of the experimental setup for neutron diffraction, i.e. it would be highly desirable to improve spatial resolution down to μ m or even sub μ m range. A straightforward approach using the same experimental geometry along with a drastic reduction of the gauge volume would lead to scattering signals below the detection limit. Therefore, new strategies and methodologies of data collection, capable to add depth resolution to the diffraction data, are highly desired, e.g. synchrotron radiation using small focused beams or developing a data collection strategy enabling inverse Radon transforms and so on.

In summary: lithium uniformity depends on different factors such as spatial current distribution, electrode morphology, electrolyte distribution and wetting, state and type of solidelectrolyte interphase, different kinds of geometrical boundary conditions *etc.*. Nature and origin of the underlying processes defining lithium distribution inside the anode are highly relevant for building safe, stable and high-performance next generation Li-ion cells. The current tabs inside the electrode rolls have been found to positively affect the cycling performance (C-rates), but cause irregularities in the electrode coating. This along with incomplete coating of the electrodes at both ends effects the shape of lithium distribution and define the islands of heterogeneity limiting the local lithium exchange.

Acknowledgements

The authors acknowledge the access to the research infrastructure of Heinz Maier-Leibnitz Zentrum (Germany) and Institut Laue – Langevin (France). The study was supported in part by German Ministry of Education and Science (Projects 05K16VK2 and 05K19VK3) and Heinz Maier-Leibnitz Zentrum. KA and FP acknowledge financial support through the DFG Gottfried Wilhelm Leibniz program. Assistance of Dr. W. Gan (Helmholtz Zentrum Geesthacht) and Dr. J. Rebello-Kornmeier (Technische Universität München) with spatially-resolved neutron diffraction experiments at STRESS-SPEC is gratefully acknowledged. We acknowledge DESY (Hamburg, Germany), a member of the Helmholtz Association HGF, for the provision of experimental facilities. Parts of this research were carried out at PETRA III using beamline P02.1. Open access funding enabled and organized by Projekt DEAL.

www.batteries-supercaps.org

334

⁶Obtained distributions of dividend and divisor in η/w quality parameter (η and 1/w) are showed vs. C-rate in Figure S7a–b along with their relationship to each other 1/w vs. η (Figure S7c).

⁷Use of high charging currents and cut-off values would potentially introduce even larger non-uniformities.



Conflict of Interest

The authors declare no conflict of interest.

Keywords: lithiated graphite · Li-ion batteries · lithium concentration · heterogeneity · neutron diffraction

- B. Scrosati, J. Hassoun, Y. K. Sun, Energy Environ. Sci. 2011, 4, 3287– 3295.
- [2] M. Armand, J. M. Tarascon, Nature 2008, 451, 652–657.
- [3] B. Diouf, R. Pode, Renewable Energy 2015, 76, 375–380.
- [4] A. Barre, B. Deguilhem, S. Grolleau, M. Gerard, F. Suard, D. Riu, J. Power Sources 2013, 241, 680–689.
- [5] V. Agubra, J. Fergus, *Materials* **2013**, *6*, 1310–1325.
- [6] J. Vetter, P. Novak, M. R. Wagner, C. Veit, K. C. Moller, J. O. Besenhard, M. Winter, M. Wohlfahrt-Mehrens, C. Vogler, A. Hammouche, J. Power Sources 2005, 147, 269–281.
- [7] D. Burow, K. Sergeeva, S. Calles, K. Schorb, A. Borger, C. Roth, P. Heitjans, J. Power Sources 2016, 307, 806–814.
- [8] P. J. Osswald, S. V. Erhard, A. Rheinfeld, B. Rieger, H. E. Hoster, A. Jossen, J. Power Sources 2016, 329, 546–552.
- [9] P. J. Osswald, S. V. Erhard, A. Noel, R. Keil, F. M. Kindermann, H. Hoster, A. Jossen, J. Power Sources 2016, 314, 93–101.
- [10] S. V. Erhard, P. J. Osswald, P. Keil, E. Hoffer, M. Haug, A. Noel, J. Wilhelm, B. Rieger, K. Schmidt, S. Kosch, F. M. Kindermann, F. Spingler, H. Kloust, T. Thoennessen, A. Rheinfeld, A. Jossen, J. Electrochem. Soc. 2017, 164, A6324–A6333.
- [11] G. S. Zhang, C. E. Shaffer, C. Y. Wang, C. D. Rahn, J. Electrochem. Soc. 2013, 160, A610–A615.
- [12] G. S. Zhang, C. E. Shaffer, C. Y. Wang, C. D. Rahna, J. Electrochem. Soc. 2013, 160, A2299–A2305.
- [13] J. Wilhelm, S. Seidlmayer, S. Erhard, M. Hofmann, R. Gilles, A. Jossen, J. Electrochem. Soc. 2018, 165, A1846–A1856.
- [14] A. Senyshyn, M. J. Muhlbauer, O. Dolotko, M. Hofmann, H. Ehrenberg, Sci. Rep. 2015, 5, 18380.
- [15] X.-L. Wang, K. An, L. Cai, Z. Feng, S. E. Nagler, C. Daniel, K. J. Rhodes, A. D. Stoica, H. D. Skorpenske, C. Liang, W. Zhang, J. Kim, Y. Qi, S. J. Harris, *Sci. Rep.* **2012**, *2*, 747.
- [16] O. Dolotko, A. Senyshyn, M. J. Muhlbauer, K. Nikolowski, H. Ehrenberg, J. Power Sources 2014, 255, 197–203.
- [17] O. Dolotko, A. Senyshyn, M. J. Muhlbauer, K. Nikolowski, F. Scheiba, H. Ehrenberg, J. Electrochem. Soc. 2012, 159, A2082–A2088.
- [18] O. Bergstrom, A. M. Andersson, K. Edstrom, T. Gustafsson, J. Appl. Crystallogr. 1998, 31, 823–825.
- [19] F. Rosciano, M. Holzapfel, W. Scheifele, P. Novak, J. Appl. Crystallogr. 2008, 41, 690–694.
- [20] M. Roberts, J. J. Biendicho, S. Hull, P. Beran, T. Gustafsson, G. Svensson, K. Edstrom, J. Power Sources 2013, 226, 249–255.
- [21] J. J. Biendicho, M. Roberts, C. Offer, D. Noreus, E. Widenkvist, R. I. Smith, G. Svensson, K. Edstrom, S. T. Norberg, S. G. Eriksson, S. Hull, J. Power Sources 2014, 248, 900–904.
- [22] N. Paul, J. Wandt, S. Seidlmayer, S. Schebesta, M. J. Muehlbauer, O. Dolotko, H. A. Gasteiger, R. Gilles, J. Power Sources 2017, 345, 85–96.
- [23] M. A. Rodriguez, M. H. Van Benthem, D. Ingersoll, S. C. Vogel, H. M. Reiche, *Powder Diffr.* 2010, 25, 143–148.
- [24] A. Senyshyn, M. J. Muhlbauer, K. Nikolowski, T. Pirling, H. Ehrenberg, J. Power Sources 2012, 203, 126–129.
- [25] A. Senyshyn, O. Dolotko, M. J. Muhlbauer, K. Nikolowski, H. Fuess, H. Ehrenberg, J. Electrochem. Soc. 2013, 160, A3198–A3205.
- [26] D. Goonetilleke, N. Sharma, W. K. Pang, V. K. Peterson, R. Petibon, J. Li, J. R. Dahn, *Chem. Mater.* **2019**, *31*, 376–386.

- [27] D. Sheptyakov, L. Boulet-Roblin, V. Pomjakushin, P. Borel, C. Tessier, C. Villevieille, J. Mater. Chem. A 2020, 8, 1288–1297.
- [28] D. Goonetilleke, J. C. Pramudita, M. Hagan, O. K. Al Bahri, W. K. Pang, V. K. Peterson, J. Groot, H. Berg, N. Sharma, J. Power Sources 2017, 343, 446–457.
- [29] S. Taminato, M. Yonemura, S. Shiotani, T. Kamiyama, S. Torii, M. Nagao, Y. Ishikawa, K. Mori, T. Fukunaga, Y. Onodera, T. Naka, M. Morishima, Y. Ukyo, D. S. Adipranoto, H. Arai, Y. Uchimoto, Z. Ogumi, K. Suzuki, M. Hirayama, R. Kanno, *Sci. Rep.* **2016**, *6*, 28843.
- [30] L. Boulet-Roblin, P. Borel, D. Sheptyakov, C. Tessier, P. Novak, C. Villevieille, J. Phys. Chem. C 2016, 120, 17268–17273.
- [31] L. Boulet-Roblin, D. Sheptyakov, P. Borel, C. Tessier, P. Novák, C. Villevieille, J. Mater. Chem. A 2017, 5, 25574–25582.
- [32] D. Petz, M. J. Mühlbauer, V. Baran, M. Frost, A. Schökel, C. Paulmann, Y. Chen, D. Garcés, A. Senyshyn, J. Power Sources 2020, 448, 227466.
- [33] M. J. Mühlbauer, D. Petz, V. Baran, O. Dolotko, M. Hofmann, R. Kostecki, A. Senyshyn, J. Power Sources 2020, 475, 228690.
- [34] V. Baran, O. Dolotko, M. J. Muhlbauer, A. Senyshyn, H. Ehrenberg, J. Electrochem. Soc. 2018, 165, A1975–A1982.
- [35] A. Senyshyn, M. J. Muhlbauer, O. Dolotko, H. Ehrenberg, J. Power Sources 2015, 282, 235–240.
- [36] L. Cai, K. An, Z. L. Feng, C. D. Liang, S. J. Harris, J. Power Sources 2013, 236, 163–168.
- [37] X. H. Yu, Z. L. Feng, Y. Ren, D. Henn, Z. G. Wu, K. An, B. Wu, C. Fau, C. Li, S. J. Harris, J. Electrochem. Soc. 2018, 165, A1578–1585.
- [38] A. Senyshyn, M. J. Muhlbauer, O. Dolotko, M. Hofmann, T. Pirling, H. Ehrenberg, J. Power Sources 2014, 245, 678–683.
- [39] M. J. Muhlbauer, O. Dolotko, M. Hofmann, H. Ehrenberg, A. Senyshyn, J. Power Sources 2017, 348, 145–149.
- [40] S. C. Nagpure, R. G. Downing, B. Bhushan, S. S. Babu, L. Cao, *Electrochim. Acta* 2011, 56, 4735–4743.
- [41] H. Kawaura, M. Harada, Y. Kondo, H. Kondo, Y. Suganuma, N. Takahashi, J. Sugiyama, Y. Seno, N. L. Yamada, ACS Appl. Mater. Interfaces 2016, 8, 9540–9544.
- [42] T. K. Zakharchenko, M. V. Avdeev, A. V. Sergeev, A. V. Chertovich, O. I. Ivankov, V. I. Petrenko, Y. Shao-Horn, L. V. Yashina, D. M. Itkis, *Nanoscale* 2019, 11, 6838–6845.
- [43] C. A. Schneider, W. S. Rasband, K. W. Eliceiri, Nat. Methods 2012, 9, 671– 675.
- [44] M. Hofmann, R. Schneider, G. A. Seidl, J. Rebelo-Kornmeier, R. C. Wimpory, U. Garbe, H. G. Brokmeier, *Phys. B* 2006, 385–86, 1035–1037.
- [45] T. Pirling, G. Bruno, P. J. Withers, *Mater. Sci. Eng. A* 2006, 437, 139–144.
- [46] C. Didier, W. K. Pang, Z. P. Guo, S. Schmid, V. K. Peterson, *Chem. Mater.* 2020, 32, 2518–2531.
- [47] N. Sharma, V. K. Peterson, M. M. Elcombe, M. Avdeev, A. J. Studer, N. Blagojevic, R. Yusoff, N. Kamarulzaman, J. Power Sources 2010, 195, 8258–8266.
- [48] J. Liu, M. Kunz, K. Chen, N. Tamura, T. J. Richardson, J. Phys. Chem. Lett. 2010, 1, 2120–2123.
- [49] D. P. Finegan, A. Quinn, D. S. Wragg, A. M. Colclasure, X. Lu, C. Tan, T. M. M. Heenan, R. Jervis, D. J. L. Brett, S. Das, T. Gao, D. A. Cogswell, M. Z. Bazant, M. Di Michiel, S. Checchia, P. R. Shearing, K. Smith, *Energy Environ. Sci.* 2020, 13, 2570–2584.
- [50] T. Kitamura, S. Takai, T. Yabutsuka, T. Yao, J. Phys. Chem. Solids 2020, 142, 109440.

Manuscript received: October 16, 2020 Revised manuscript received: October 26, 2020 Accepted manuscript online: October 28, 2020 Version of record online: November 23, 2020

335